Schreiber, David

From:

Lacourciere, Karen

Sent:

Monday, September 29, 2003 6:45 PM

To:

Schreiber, David

Subject:

Specialized Sequence Search request 09/676,436

Hi David Schreiber-

I was wondering if you could run the type of specialized search we discussed for antisense sequences, wherein you rank ordered the hits using an Excel spreadsheet to pull up short, hit percent match hits for 09/676,436?

Could you please run a <u>length limited</u> nucleotide sequence search against nucleotides 3314-3333 of SEQ ID NO:3 that returns hits 8-50 nucleotides then rank orders the hits based on percent complementarity over the whole oligo, exactly as you performed previously? Please call me if there are any questions. The number of hits to collect in the first round would be your best judgement.

Thanks, Karen

* Karen A. Lacourciere Ph. D. CM1 11D09 GAU 1635 (703) 308-7523

Access		

SEARCH REQUEST FORM

Scientific and Technical Information Center

Requester's Full Name		Examiner # :Serial Number:	Date:
Art Unit Phone	Number 30	Serial Number:	DADED DION ENAM
Mail Box and Bldg Room Location	on Kc	sults Format Preferred (circle)	PAPER DISK E-MAIL
If more than one search is sub	mitted, please priorit		eed. **************
Please provide a detailed statement of the Include the elected species or structures utility of the invention. Define any term known Please attach a copy of the cove	, keywords, synonyms, acro is that may have a special r	onyms, and registry numbers, and meaning. Give examples or releva	combine with the concept or
Title of Invention			0
Inventors (please provide full names):			
		:	A Caracana and a cara
Earliest Priority Filing Date.			"
*For Sequence Searches Only * Please inc appropriate social number.			patent numbers) along with the
STAFF USE ONLY Searcher	Type of Search NA Sequence (#)		where applicable
Searcher Phone =			
Searcher Location			
Dare Nearther Filikoli in			
Dare impleted		.evs Nevs	
Searcher Pred & Religial Time [1]	Full text	Nequente Rystems	
Clenical Prepiir mit	Patent Family	semetral X-X-X	
	Other	Other specify	

GenCore version 5.1.6 Copyright (c) 1993 -

2003 Compugen Ltd.

OM nucleic - nucleic using sw model

Run on: Septem 30, 2003, 7:46:12;

(without alignments)

199.182 Million cell updates/sec

Title: US-09-676-436-3_COPY_3314_3333

Perfect score: 20

Sequenc 1 agggattcaggggttccagc 20

Scoring table: IDENTITY_NUC

Gapop 10 , Gapex 1

Searched 33363688 seqs, 1.7E+10 residues

Total number of hits satisfy chosen parameters 21323498

MinimumDB seq length: 8
MaximumDB seq length: 50

Post-prodMinimum Match 0% Maximum Match 100%

Listing first 65000 summaries

Database Pending Patents NA Main:*

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Pred.	No.	is	the	number	of	results	predicted
score	greater	than	or	equal	to	the	score
and	is	derived	by	analysis	of	the	total

SUMMARIES

%

Result No.	Score		Query Match	Length	DB	ID	Score/Length
c 1		20	100	20		1 PCT-US01-30549-12	1
c 2		20	100	20		2 PCT-US01-30549-12	1
c 3		20	100	20	2	9 US-09-676-436-12	1
c 4		20	100	20	5	51 US-10-371-474-12	1
c18928		10	50	10	. 1	9 US-09-336-376-2856	1

c18929	10	50	10	19 US-09-336-376-4653	1
517	12.4	62	14	10 US-08-591-486B-72	0.885714
518	12.4	62	14	19 US-09-341-700A-895	0.885714
c41067	9.4	47	11	50 US-10-320-210A-4	0.854545
c41068	9.4	47	11	50 US-10-320-210A-127	0.854545
c18930	10	50	12	22 US-09-528-209A-8913	0.833333

OM nucleic - nucleic search, using sw model

Run on:

September 30, 2003, 08:22:42; Search time 1924 Seconds

(without alignments)

25.878 Million cell updates/sec

Title:

US-09-676-436-3 COPY_3314_3333

Perfect score: 20

Sequence:

1 agggattcaggggttccagc 20

Scoring table: IDENTITY NUC

Gapop 10.0, Gapext 1.0

Searched:

1678620 segs, 1244745471 residues

Total number of hits satisfying chosen parameters:

1072644

Minimum DB seq length: 8 Maximum DB seq length: 50

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 65000 summaries

Database:

Published Applications NA:*

- 1: /cgn2_6/ptodata/2/pubpna/US07_PUBCOMB.seq:*
- 2: /cgn2_6/ptodata/2/pubpna/PCT_NEW_PUB.seq:*
- 3: /cgn2 6/ptodata/2/pubpna/US06 NEW PUB.seg:*
- 4: /cgn2_6/ptodata/2/pubpna/US06 PUBCOMB.seq:*
- 5: /cgn2 6/ptodata/2/pubpna/US07 NEW PUB.seq:*
- 6: /cgn2 6/ptodata/2/pubpna/PCTUS PUBCOMB.seg:*
- 7: /cgn2 6/ptodata/2/pubpna/US08 NEW PUB.seq:*
- 8: /cgn2_6/ptodata/2/pubpna/US08_PUBCOMB.seq:*
- 9: /cgn2_6/ptodata/2/pubpna/US09A_PUBCOMB.seq:*
- 10: /cgn2 6/ptodata/2/pubpna/US09B PUBCOMB.seq:*
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- 13: /cgn2 6/ptodata/2/pubpna/US10A PUBCOMB.seq:*
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- 17: /cgn2 6/ptodata/2/pubpna/US60_PUBCOMB.seq:*

Pred. No. is the number of results predicted by chance to have a

score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

%	
70	

Result No.	Score	Query Match Ler	ngth DB	ID	Score/length
c 1	20	100	20	12 US-10-371-474-12	1
22	12.4	62	14	8 US-08-591-486B-72	0.885714
8396	8.4	42	10	9 US-09-989-789-1270	0.84
8397	8.4	42	10	9 US-09-989-789-1275	0.84
8398	8.4	42	10	9 US-09-989-789-1337	0.84
8399	8.4	42	10	11 US-09-990-186-1270	0.84
8400	8.4	42	10	11 US-09-990-186-1275	0.84
8401	8.4	42	10	11 US-09-990-186-1337	0.84
c8402	8.4	42	10	11 US-09-953-562-26	0.84
8403	8.4	42	10	11 US-09-989-994-1270	0.84
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8406	8.4	42	10	12 US-10-330-627-435	0.84
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8408	8.4	42	10	12 US-10-330-627-1546	0.84
c29527	7.4	37	9	9 US-09-955-518-18	0.822222
14156	8	40	10	12 US-10-330-627-588	0.8
c14157	8	40	10	13 US-10-033-145-524	8.0
14158	8	40	10	13 US-10-033-145-765	8.0
14159	8	40	10	13 US-10-033-145-1337	0.8

OM nucleic - nucleic search, using sw mode!

Run on:

September 30, 2003, 07:46:12; Search time 283 Seconds

(without alignments)

190.773 Million cell updates/sec

Title:

US-09-676-436-3_COPY_3314_3333

Perfect score: 20

Sequence:

1 agggattcaggggttccagc 20

Scoring table: IDENTITY NUC

Gapop 10.0, Gapext 1.0

Searched:

2552756 segs, 1349719017 residues

Total number of hits satisfying chosen parameters:

2466760

Minimum DB seq length: 8
Maximum DB seq length: 50

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 65000 summaries

Database:

N Geneseg 19Jun03:*

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- 6: /SIDS1/gcgdata/geneseq/geneseqn-embl/NA1985.DAT:*
- 7: /SIDS1/gcgdata/geneseq/geneseqn-embl/NA1986.DAT:*
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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

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			enath	DB II	D	Score/Length
						_
c 1	20	100	20	24	AAD37150	1
c 427	11	55	11	24	ABV67711	1
c 428	11	55	11	24	ABQ87435	1
c1844	10	50	10	21	AAZ83859	1
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c1845	10	50	11		ABV66038	
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6906	9	45	10		AAF42053	
c6907	9	45	10		AAL48142	
c6908	9	45			ABK64082	
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4117	9.4	47	11		ABV68833	
c4118	9.4	47	11		ABV70104	
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c14568	8.4	42	10	_	AAZ83550	0.84
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c14570	8.4	42			AAZ84773	0.84
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c14573	8.4				AAZ85716	0.84
c14574	8.4				: AAI67389	0.84
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c14580	8.4	42	10	22 AAH19941	0.84
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c14582	8.4	42	10	22 AAH32697	0.84
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24575	8	40	10	21 AAZ84157	0.8
c24576	8	40	10	21 AAZ84158	0.8
c24577	8	40	10	21 AAZ84257	0.8
c24578	8	40	10	21 AAZ84401	0.8
24579	8	40	10	21 AAZ84685	0.8
24580	8	40	10	21 AAZ85131	0.8
c24581	8	40	10	21 AAZ85236	0.8
24582	8	40	10	22 AAH63748	0.8
24583	8	40	10	22 AAF35691	0.8
c24584	8	40	10	22 AAF36466	0.8
c24585	8	40	10	22 AAF40055	0.8
24586	8	40	10	22 AAF42054	0.8
24587	8	40	10	22 AAF42057	0.8
24588	8	40	10	22 AAF42631	6.0
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c24589	8	40	10	22 AAF42841	0.8
24590	8	40	10	24 AAD45882	0.8
c24591	8	40	10	24 ABK96539	8.0
c24592	8	40	10	24 ABK92637	8.0
24593	8	40	10	24 ABK14251	8.0
24594	8	40	10	24 ABK23538	0.8
24595	8	40	10	24 ABK23629	0.8
c24596	8	40	10	25 ABT14312	0.8

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OM nucleic - nucleic search, using sw model

Run on:

September 30, 2003, 07:46:12; Search time 1801 Seconds

(without alignments)

454.299 Million cell updates/sec

Title:

US-09-676-436-3_COPY_3314_3333

Perfect score: 20

Sequence:

1 agggattcaggggttccagc 20

Scoring table: IDENTITY NUC

Gapop 10.0, Gapext 1.0

Searched:

2888711 segs, 20454813386 residues

Total number of hits satisfying chosen parameters:

1314098

Minimum DB seq length: 8 Maximum DB seq length: 50

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 65000 summaries

Database:

GenEmbl:*

- 1: gb_ba:*
- 2: gb_htg:*
- 3: gb_in:*
- 4: gb_om:*
- 5: gb_ov:*
- 6: gb_pat:*
- 7: gb_ph:*
- 8: gb pl:*
- 9: gb_pr:*
- 10: gb_ro:*
- 11: gb_sts:*
- 12: gb_sy:*
- 13: gb_un:*
- 14: gb_vi:*
- 15: em ba:*
- 16: em_fun:*
- 17: em hum:*
- 18: em_in:*
- 19: em_mu:*

20: em_om:* 21: em_or:* 22: em_ov:* 23: em_pat:* 24: em_ph:* 25: em_pl:* 26: em_ro:* 27: em_sts:* 28: em_un:* 29: em_vi:* 30: em_htg_hum:* 31: em_htg_inv:* 32: em_htg_other:* 33: em_htg_mus:* 34: em_htg_pln:* 35: em_htg_rod:* 36: em_htg_mam:* 37: em_htg_vrt:* 38: em_sy:* 39: em_htgo_hum:* 40: em_htgo_mus:* 41: em_htgo_other:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

		%				
Result		Query				
No.	Score	Match	Length	DB I	D	Score/length
c 329	11	55	5 11	6	AX471613	1
c 330	11	55	5 11	6	AX628456	1
c1489	10	50) 10) 6	184475	1
c1490	10	50) 11	6	AX626783	0.909090909
5458	9	45	5 10) 6	E39766	0.9
36	12.4	62	! 14	6	A42556	0.885714286
37	12.4	62	2 14	6	A88747	0.885714286
38	12.4	62	2 14	6	BD066260	0.885714286
c3262	9.4	47	' 11	6	AX470549	0.854545455
c3263	9.4	47	' 11	6	AX623428	0.854545455
3264	9.4	47	7 11	6	AX629578	0.854545455
c3265	9.4	47	' 11	1 6	AX:630849	0.854545455
11538	8.4	42	2 10	6 (AR303398	0.84
c11539	8.4	42	2 10	0 6	AR303424	0.84
c11540	8.4	42	2 10) 6	AX112965	0.84
11541	8.4	42	2 10) 6	AX152520	0.84
11542	8.4	42	2 10) 6	AX153581	0.84

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8.4	42	10	6 AX153631	0.84
8.4	42	10	6 AX301326	0.84
8.4	42	10	6 AX667821	0.84
8.4	42	10	6 AX667826	0.84
8.4	42	10	6 AX667888	0.84
8.4	42	10	6 BD007762	0.84
8.4	42	10	6 BD007794	0.84
8.4	42	10	6 BD083142	0.84
8.4	42	10	6 BD083204	0.84
8.4	42	10	6 BD083268	0.84
8.4	42	10	6 E39536	0.84
8.4	42	10	6 E54753	0.84
9	45	11	6 AX471317	0.818181818
9	45	11	6 AX624360	0.818181818
9	45	11	6 AX625683	0.818131818
9	45	11	6 AX625706	0.818181818
9	45	11	6 AX626201	0.818181818
9	45	11	6 AX631781	0.818181818
8	40	10	6 AR017955	8.0
8	40	10	6 AR303474	8.0
8	40	10	6 AR303528	0.8
8	40	10	6 AX152673	8.0
8	40	10	6 AX301451	8.0
8	40	10	6 AX301542	0.8
8	40	10	6 AX719148	8.0
	8.4 8.4 8.4 8.4 8.4 8.4 8.4 9 9 9 9 9 9 9 9 8 8 8 8 8 8 8 8 8 8 8	8.4 42 8.4 42 8.5 9 45 9 45 8 40 8 80 8 80 8 80 8 80 8 80 8 80 8 80 8 80 8 80 8 80	8.4 42 10 8.4 42 10 8.4 42 10 8.4 42 10 8.4 42 10 8.4 42 10 8.4 42 10 8.4 42 10 8.4 42 10 8.4 42 10 9 45 11 9 45 11 9 45 11 9 45 11 9 45 11 9 45 11 9 45 11 9 45 11 9 45 11 9 45 11 9 45 11 9 45 11 9 45 11 9 45 11 9 45 10 8 40 10 8 40 10 8 40 10 <td>8.4 42 10 6 AX301326 8.4 42 10 6 AX667821 8.4 42 10 6 AX667826 8.4 42 10 6 BD007762 8.4 42 10 6 BD007794 8.4 42 10 6 BD083142 8.4 42 10 6 BD083204 8.4 42 10 6 BD083204 8.4 42 10 6 BD083204 8.4 42 10 6 BD083268 8.4 42 10 6 E39536 8.4 42 10 6 E54753 9 45 11 6 AX471317 9 45 11 6 AX624360 9 45 11 6 AX625683 9 45 11 6 AX625706 9 45 11 6 AX625706 9 45 11 6 AX625706 9 45 11 6 AX626201 9 45 11 6 AX631781 8 40 10 6 AR303474 8 40 10 6 AR303528 8 40 10 6 AX301451 8 40 10 6 AX301451 8 40 10 6 AX301542</td>	8.4 42 10 6 AX301326 8.4 42 10 6 AX667821 8.4 42 10 6 AX667826 8.4 42 10 6 BD007762 8.4 42 10 6 BD007794 8.4 42 10 6 BD083142 8.4 42 10 6 BD083204 8.4 42 10 6 BD083204 8.4 42 10 6 BD083204 8.4 42 10 6 BD083268 8.4 42 10 6 E39536 8.4 42 10 6 E54753 9 45 11 6 AX471317 9 45 11 6 AX624360 9 45 11 6 AX625683 9 45 11 6 AX625706 9 45 11 6 AX625706 9 45 11 6 AX625706 9 45 11 6 AX626201 9 45 11 6 AX631781 8 40 10 6 AR303474 8 40 10 6 AR303528 8 40 10 6 AX301451 8 40 10 6 AX301451 8 40 10 6 AX301542

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OM nucleic - nucleic search, using sw model

Run on:

September 30, 2003, 07:46:12; Search time 74 Seconds

(without alignments)

119.293 Million cell updates/sec

Title:

US-09-676-436-3_COPY_3314_3333

Perfect score: 20

Sequence:

1 agggattcaggggttccagc 20

Scoring table: IDENTITY_NUC

Gapop 10.0, Gapext 1.0

Searched:

569978 segs, 220691566 residues

Total number of hits satisfying chosen parameters:

741060

Minimum DB seg length: 8 Maximum DB seq length: 50

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 65000 summaries

Database:

Issued_Patents_NA:*

- 1: /cgn2 6/ptodata/2/ina/5A COMB.seq:*
- 2: /cgn2 6/ptodata/2/ina/5B COMB.seq:*
- 3: /cgn2 6/ptodata/2/ina/6A COMB.seg:*
- 4: /cgn2_6/ptodata/2/ina/6B_COMB.seq:*
- 5: /cgn2_6/ptodata/2/ina/PCTUS_COMB.seq:*
- 6: /cgn2_6/ptod.ata/2/ina/backfiles1.seq:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

%

Result No.	Query Score Match	Length	DB	ID	Score/Length
c 955	10	50 1	0	1 US-07-704-288C-25	1
c 956	10	50 1	0	1 US-08-379-259-25	1

c11470	8	40	8	3 US-08-859-954-239	1
37837	7	35	8	3 US-08-859-954-32	0.875
c37838	7	35	8	3 US-08-859-954-240	0.875
c37839	7	35	8	3 US-08-859-954-292	0.875
c37840	7	35	8	3 US-09-063-450-8	0.875
c37841	7	35	8	3 US-09-398-499-15	0.875
37842	7	35	8	3 US-09-398-499-38	0.875
7013	8.4	42	10	4 US-09-508-753B-123	0.84
c7014	8.4	42	10	4 US-09-508-753B-149	0.84
c23020	7.4	37	9	2 US-08-899 - 324-6	0.822222
c23021	7.4	37	9	3 US-08-329-892B-6	0.822222
c23022	7.4	37	9	4 US-08-874-569B-18	0.822222
c23023	7.4	37	9	4 US-09-955-518-18	0.822222
11471	8	40	10	1 US-08-590-804-20	0.8
11472	8	40	10	4 US-09-508-753B-199	0.8
c11473	8	40	10	4 US-09-508-753B-253	0.8